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#### **PCT**

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An acidic phospholipase is obtained from a strain of the genus *Hyphozyma*. It is able to hydrolyze both fatty acyl groups in intact phospholipid. Advantageously, it has no lipase activity and is active at very low pH; these properties make it very suitable for use in oil degumming, as enzymatic and alkaline hydrolysis (saponification) of the oil can both be suppressed. The phospholipase is not membrane bound, making it suitable for commercial production and purification.

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#### NOVEL PHOSPHOLIPASE, PRODUCTION AND USE THEREOF

#### TECHNICAL FIELD

This invention relates to a novel phospholipase, DNA encoding it and to its production and use.

#### 5 BACKGROUND ART

Phospholipids, such as lecithin or phosphatidylcholine, consist of glycerol esterified with two fatty acids in an outer (sn-1) and the middle (sn-2) positions and esterified with phosphoric acid in the third position; the phosphoric acid, in turn, may be esterified to an amino-alcohol. Phospholipases are enzymes which participate in the hydrolysis of phospholipids. Several types of phospholipase activity can be distinguished, including phospholipase A1 and A2 which hydrolyze one fatty acyl group (in the sn-1 and sn-2 position, respectively) to form lysophospholipid; and lysophospholipase (or phospholipase B) which can hydrolyze the remaining fatty acyl group in lysophospholipid. This invention relates to a phospholipase that has the ability to hydrolyze both fatty acyl groups in a phospholipid. Enzymes with this activity are also some times called phospholipase B.

Enzymes with phospholipase B activity have been reported from various fungal sources, including *Penicillium notatum* (also known as *P. chrysogenum*; N. Kawasaki, J. Biochem., 77, 1233-44, 1975; N. Masuda et al., Eur. J. Biochem., 202, 783-787, 1991), *Saccharomyces cerevisiae* (M. Ichimasa et al., Agric. Biol. Chem., 49 (4), 1083-89, 1985; F. Paultauf et al., J. Biol. Chem., 269, 19725-30, 1994), *Torulaspora delbrueckii* (old name *Saccharomyces rosei*; Y. Kuwabara, Agric. Biol. Chem., 52 (10), 2451-58, 1988; FEMS, Microbiol. Letters, 124, 29-34), *Schizosaccharomyces pombe* (H. Oishi et al., Biosci. Biotech. Biochem., 60 (7), 1087-92, 1996), *Aspergillus niger* (Technical Bulletin, G-zyme™ G999, Enzyme Bio-Systems Ltd.) and *Corticium centrifugum* (S. Uehara et al., Agric. Biol. Chem., 43 (3), 517-525, 1979).

It is known to use phospholipase in, e.g., enzymatic oil degumming (US 5,264,367, Metallgesellschaft, Röhm), treatment of starch hydrolysate (particularly from wheat starch) to improve the filterability (EP 219,269, CPC International) and as an additive to bread dough to improve the elasticity of the bread (US 4,567,046, Kyowa Hakko).

It is the object of this invention to provide an improved phospholipase for use in such processes.

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#### STATEMENT OF THE INVENTION

The present inventors have found that an acidic phospholipase can be obtained from a strain of the genus Hyphozyma. It is able to hydrolyze both fatty acyl groups in intact phospholipid. Advantageously, it has no lipase activity and is active at 5 very low pH; these properties make it very suitable for use in oil degumming, as enzymatic and alkaline hydrolysis (saponification) of the oil can both be suppressed. The phospholipase is not membrane bound, making it suitable for commercial production and purification.

WO 93/24619 (Novo Nordisk) discloses a lipase from Hyphozyma sp. LF-132 10 (CBS 648.91), but the production of phospholipase by this genus has never been reported. We have found that the phospholipase of this invention can be obtained from the same strain as the known lipase, and that the two enzymes can be separated.

Accordingly, a first aspect of the invention provides an isolated phospholipase which is able to hydrolyze both fatty acyl groups in a phospholipid, is derivable from a 15 strain of Hyphozyma, and has optimum phospholipase activity at about 50°C and pH 3 measured at the conditions described in Example 3.

The invention also provides an isolated phospholipase which is able to hydrolyze both fatty acyl groups in a phospholipid, and is a polypeptide comprising at its N-terminal a partial amino acid sequence which is the sequence shown in positions 20 1-497 of SEQ ID NO: 11, or is at least 50 % identical therewith

In another aspect, the invention provides an isolated phospholipase which is able to hydrolyze both fatty acyl groups in a phospholipid, and is a polypeptide containing amino acid sequences which are at least 50 % identical with the amino acid sequences shown in SEQ ID NO: 1-8, disregarding Xaa.

The invention further provides an isolated DNA sequence which encodes said phospholipase.

Yet another aspect of the invention provides a method of producing a phospholipase, comprising cultivation of a phospholipase-producing strain of Hyphozyma in a suitable nutrient medium, followed by recovery of the phospholipase

A further aspect of the invention provides a method for producing a phospholipase, comprising isolating a DNA sequence encoding the phospholipase from a phospholipase-producing strain of Hyphozyma, combining the DNA fragment with appropriate expression signal(s) in an appropriate vector, transforming a suitable heterologous host organism with the vector, cultivating the transformed host organism 35 under conditions leading to expression of the phospholipase, and recovering the phospholipase from the culture medium

The invention also provides use of said phospholipase in a process comprising treatment of a phospholipid or lysophospholipid with the phospholipase so as to hydrolyze fatty acyl groups.

Finally, the invention provides a process for reducing the content of phospholipid in a vegetable oil, comprising treating the oil with an aqueous dispersion of an acidic phospholipase at pH 1.5-3 so as to hydrolyze a major part of the phospholipid, and separating an aqueous phase containing the hydrolyzed phospholipid from the oil.

#### **BRIEF DESCRIPTION OF DRAWINGS**

Figs. 1, 2 and 3 show the temperature profile, pH profile and thermostability, respectively, of phospholipase from *Hyphozyma sp.* CBS 648.91. Further details are given in Example 3.

Fig. 4a-d gives a comparison of SEQ ID NO: 11 with 3 prior-art sequences.

#### DETAILED DISCLOSURE OF THE INVENTION

#### 15 Phospholipase

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The phospholipase of the invention is able to hydrolyze both acyl groups in a phospholipid molecule (such as phosphatidyl choline or lecithin) without intermediate accumulation of lysophospholipid and is also able to hydrolyze the fatty acyl group of a lysophospholipid (such as lysophosphatidyl choline or lyso-lecithin). Advantageously, the phospholipase of the invention is not membrane bound.

A preferred enzyme is derived from *Hyphozyma sp.* strain CBS 648.91. Its molecular weight is about 94 kDa by SDS, about 87 kDa by gel filtration, and 92 kDa by mass spectrometry. It is believed to be glycosylated. It has an iso-electric point of about 5.6. It has no lipase activity, i.e. it does not hydrolyze triglycerides.

The influence of pH and temperature on the activity of this phospholipase is shown in Fig. 1 and 2. As shown in these figures, the enzyme has optimum activity at about pH 3 and 50°C.

Fig. 3 shows the thermostability of this enzyme, expressed as the residual activity after 10 minutes at pH 7 at various temperatures. It is seen that the enzyme retains more than 90 % activity at temperatures up to 50°C, more than 75% up to 60°C and more than 50% up to 70°C.

#### **Phospholipase Activity Assay**

Two different units are used in this specification:

1 unit (phospholipase activity unit) is the amount of phospholipase that releases one  $\mu(\text{micro})$ -mole of fatty acid per minute from DPPC (dipalmitoyl phosphatidylcholine) at 40°C and pH 4. The amount of released fatty acid is determined by NEFA-C test Wako.

1 International Unit (IU) is the amount of phospholipase that releases one  $\mu(\text{micro})$ -equivalent of free fatty acid per minute from egg yolk in the presence of calcium and deoxycholate at pH 8.0 and 40°C in a pH-stat. The released fatty acids are titrated with 0.1 N sodium hydroxide and the base volume is monitored as a function of time.

#### 10 Assay for action pattern of phospholipase

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The following test is used to identify if a given enzyme has the ability to hydrolyze both fatty acyl groups of a phospholipid without the accumulation of lysophospholipid.

A substrate solution is prepared containing 2% L-α(alpha)15 phosphatidylcholine, dipalmitoyl (product of Wako Pure Chemical Industries Ltd.) and
2% Triton X-100. A buffer solution is prepared containing 0.4 M citrate buffer (pH 5).
Enzyme solutions are prepared containing various amounts of the sample to be analyzed.

0.5 ml of the substrate solution, 0.25 ml of the buffer solution and 0.05 ml of 20 0.1 N CaCl<sub>2</sub> are mixed and incubated at 40°C. 0.1 ml of the enzyme solution is added and incubated for 1 hour. The reaction is terminated by adding 0.1 ml of 1 N HCl.

2 ml of CHCl<sub>3</sub>-methanol (1:1) is added to the reaction mixture and mixed vigorously. Approx. 1 μ(micro)l of the CHCl<sub>3</sub>-methanol is taken and applied to a TLC rod (in triplicate or quadruplicate). the TLC rods are dried and developed for 45 minutes with CHCl<sub>3</sub>: methanol: NH<sub>3</sub> (25% solution) = 65:25:5. After the development, the rods are scanned by TLC-FID (latroscan), and the chromatograms are integrated.

The amounts of palmitate, the substrate, lysophosphatidyl choline (LPC) and glycerophosphatidyl choline (GPC) are calculated from the areas of peaks appearing in that order.

The result of the test is considered positive if GPC is formed without any LPC formation.

#### Amino acid sequence

Partial sequences SEQ ID NO: 1-8 were determined by sequencing of phospholipase from *Hyphozyma sp.* CBS 648.91 after enzymatic hydrolysis. In these sequences, Xaa represents an amino acid that could not be determined. SEQ ID NO:

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1 is an N-terminal sequence, and the others are internal sequences. Xaa in SEQ ID NO: 1 is believed to be a Pro residue. Xaa in SEQ ID NO: 3, 7 and 8 and both Xaa in SEQ ID NO: 5 are believed to be glycosylated Asn residues.

A nearly complete DNA sequence (SEQ ID NO: 9) was determined for the 5 gene encoding the phospholipase from Hyphozyma sp. CBS 648.91. This sequence was determined from the genomic locus and includes an open reading frame of 552 amino acids and 213 base pairs of sequence upstream of the putative translation initiation codon. The methods used for sequence isolation and determination are well known in the art. Details are given in the examples.

The long, uninterrupted open reading frame identified in this sequence was translated and compared to the partial peptide sequences SEQ ID NO: 1-8. The translated sequence was identical to seven of the partial peptide sequences at all positions, SEQ ID NO:1-7, and overlapped the most distal partial peptide sequence. SEQ ID NO: 8 by 10 amino acids. By combining the translation with partial peptide 15 NO: 8, a sequence of 573 amino acid residues (shown as SEQ ID NO: 11) has been determined. The amino terminus of the mature peptide is determined by comparison with SEQ ID NO: 1. The sequenced open reading frame extends upstream an additional 115 amino acids. There is only one Met codon in this region, 76 amino acids from the start of the mature peptide (position -76). The 14 amino acids immediately 20 following this methionine residue appear to constitute a secretion signal sequence (G. von Heijne, Nucleic Acids Res, 14, 4683-4690, 1986), indicating both that this is the translation initiation codon and that the encoded protein is secreted. The intervening 61 amino acids must constitute a propeptide.

The peptide sequence from Hyphozyma was aligned with the phospholipase B 25 sequences from three other fungi, Penicillium notatum (Genbank X60348). Saccharomyces cerevisiae (Genbank L23089) and Torulaspora delbrueckii (Genbank D32134), as shown in Fig. 4a-d. In this alignment a dash (-) indicates an inserted gap, a circle (o) above the alignment marks a position at which the same amino acid is found in all proteins, and a vertical line (I) above the alignment indicates similar 30 residues in all proteins. The portion of the Hyphozyma phospholipase sequence we have determined is 38% identical to the phospholipase from Penicillium notatum, 37% identical to the phospholipase from Saccharomyces cerevisiae, and 38% identical to the phospholipase from Torulaspora delbrueckii. The full length Penicillium, Saccharomyces, and Torulaspora sequences extend from 112 to 145 residues further 35 than the partial Hyphozyma sequence, suggesting that the full length for the translated Hyphozyma peptide is approximately 700 amino acid residues.

Thus, the phospholipase of the invention may contain an N-terminal sequence as shown at positions 1-497 of SEQ ID NO: 11 or a sequence derived therefrom by substitution, deletion or insertion of one or more amino acids. The derived sequence may be at least 50 % identical, e.g. at least 60%, preferably at least 70%, especially at least 80 or at least 90% identical with said partial sequence. The phospholipase of the invention may contain a further 150-250 (e.g. 180-220) amino acid residues at the C-terminal

#### Microorganism

The phospholipase of this invention may be derived from a fungal strain of the genus *Hyphozyma*, a genus of yeast-like *Hyphomycetes* described in de Hoog, G.S & Smith, M.Th., Antonie van Leeuwenhoek, 47, 339-352 (1981).

Preferably, the strain belongs to the species defined by the strain *Hyphozyma sp.* LF132, CBS 648.91, which is described in WO 93/24619. This strain was classified in the genus Hyphozyma, but it did not match any of the previously described species of *Hyphozyma*, so it is believed to define a new species. It is particularly preferred to use said strain or a mutant or variant thereof having the ability to produce phospholipase.

The preferred *Hyphozyma sp.* strain (designated LF132 by the inventors) has been deposited on 12 November 1991, for the purpose of patent procedures according to the Budapest Treaty at Centraal Bureau voor Schimmelcultures (CBS), Oosterstraat 1, 3740 AG Baarn, Netherlands, and was given the accession number CBS 648.91.

#### Production of phospholipase by cultivation of Hyphozyma

The phospholipase of the invention may be produced by cultivation of the microorganism described above in a suitable nutrient medium, containing carbon and nitrogen sources and inorganic salts, followed by recovery of the enzyme. The nutrient medium may be formulated according to principles well known in the art.

The phospholipase may be recovered from the culture broth and purified to remove lipase activity, e.g. as described in the examples of this specification.

#### 30 Production by cultivation of transformant

An alternative method of producing the phospholipase of the invention comprises transforming a suitable host cell with a DNA sequence encoding the phospholipase, cultivating the transformed organism under conditions permitting the production of the enzyme, and recovering the enzyme from the culture.

The host organism is preferably a eukaryotic cell, in particular a fungal cell, such as a yeast cell or a filamentous fungal cell, preferably a strain of Aspergillus, Fusarium, Trichoderma or Saccharomyces, most preferably A. niger, A. oryzae, F. graminearum, F. sambucinum, F. cerealis or S. cerevisiae. The production of the phospholipase in such host organisms may be done by the general methods described in EP 238,023 (Novo Nordisk), WO 96/00787 (Novo Nordisk) or EP 244,234 (Alko).

The DNA sequence can be isolated from a phospholipase-producing *Hyphozyma* strain by extraction of DNA by methods known in the art, e.g. as described by Sambrook et al., (1989), Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Lab.; Cold Spring Harbor, NY.

The DNA sequence of the invention can also be isolated by any general method involving

- cloning, in suitable vectors, a cDNA library from a phospholipase-producing 15 *Hyphozyma* strain,
  - transforming suitable yeast host cells with said vectors.
  - culturing the host cells under suitable conditions to express any enzyme of interest encoded by a clone in the cDNA library,
- screening for positive clones by determining any phospholipase activity of the enzyme produced by such clones, and
  - isolating the enzyme encoding DNA from such clones.

A general isolation method has been disclosed in WO 93/11249 or WO 94/14953, the contents of which are hereby incorporated by reference.

Alternatively, the DNA encoding a phospholipase of the invention may, in accordance with well-known procedures, conveniently be isolated from a phospholipase-producing *Hyphozyma* strain, by use of synthetic oligonucleotide probes prepared on the basis of a peptide sequence disclosed herein.

#### Use of phospholipase

The phospholipase of the invention can be used in any application where it is desired to hydrolyze the fatty acyl group(s) of a phospholipid or lyso-phospholipid, such as lecithin or lyso-lecithin. The phospholipase is preferably used at pH 1.5-5 (e.g. 3-5, particularly 3.5-4.5) and at 30-70°C (particularly 40-60°C). If desired, the phospholipase may be inactivated after the reaction by a heat treatment, e.g. at pH 7, 80°C for 1 hour or 90°C for 10 minutes.

As an example, the phospholipase of the invention can be used in the preparation of dough, bread and cakes, e.g. to improve the elasticity of the bread or

cake. Thus, the phospholipase can be used in a process for making bread, comprising adding the phospholipase to the ingredients of a dough, kneading the dough and baking the dough to make the bread. This can be done in analogy with US 4,567,046 (Kyowa Hakko), JP-A 60-78529 (QP Corp.), JP-A 62-111629 (QP Corp.), JP-A 63-5 258528 (QP Corp.) or EP 426211 (Unilever).

The phospholipase of the invention can also be used to improve the filterability of an aqueous solution or slurry of carbohydrate origin by treating it with the phospholipase. This is particularly applicable to a solution or slurry containing a starch hydrolysate, especially a wheat starch hydrolysate since this tends to be difficult to filter and to give cloudy filtrates. The treatment can be done in analogy with EP 219,269 (CPC International).

#### Treatment of vegetable oil

The phospholipase of the invention can be used in a process for reducing the content of phospholipid in an edible oil, comprising treating the oil with the phospholipase so as to hydrolyze a major part of the phospholipid, and separating an aqueous phase containing the hydrolyzed phospholipid from the oil. This process is applicable to the purification of any edible oil which contains phospholipid, e.g. vegetable oil such as soy bean oil, rape seed oil and sunflower oil.

Prior to the enzymatic treatment, the vegetable oil is preferably pretreated to remove slime (mucilage), e.g. by wet refining. Typically, the oil will contain 50-250 ppm of phosphorus as phospholipid at the start of the treatment with phospholipase, and the process of the invention can reduce this value to below 5-10 ppm.

The enzymatic treatment is conducted by dispersing an aqueous solution of the phospholipase, preferably as droplets with an average diameter below 10 µ(micro)m. The amount of water is preferably 0.5-5% by weight in relation to the oil. An emulsifier may optionally be added. Mechanical agitation may be applied to maintain the emulsion.

The enzymatic treatment can be conducted at a pH in the range 1.5-5. The process pH may be in the range 3.5-5 in order to maximize the enzyme performance, or a pH in the range 1.5-3 (e.g. 2-3) may be used in order to suppress the alkaline hydrolysis of triglycerides (saponification). The pH may be adjusted by adding citric acid, a citrate buffer or HCI.

A suitable temperature is generally 30-70°C (particularly 30-45°C, e.g. 35-40°C). The reaction time will typically be 1-12 hours (e.g. 2-6 hours), and a suitable enzyme dosage will usually be 100-5000 IU per liter of oil (e.g. 200-2000 IU/I) or 0.1-10 mg/l (e.g. 0.5-5 mg/l).

The enzymatic treatment may be conducted batchwise, e.g. in a tank with stirring, or it may be continuous, e.g. a series of stirred tank reactors.

The enzymatic treatment is followed by separation of an aqueous phase and an oil phase. This separation may be performed by conventional means, e.g. 5 centrifugation. The aqueous phase will contain phospholipase, and the enzyme may be re-used to improve the process economy.

In other respects, the process can be conducted according to principles known in the art, e.g. in analogy with US 5,264,367 (Metallgesellschaft, Röhm); K. Dahlke & H. Buchold, INFORM, 6 (12), 1284-91 (1995); H. Buchold, Fat Sci. Technol., 95 (8), 300-304 (1993); JP-A 2-153997 (Showa Sangyo); or EP 654,527 (Metallgesellschaft, Röhm).

#### **EXAMPLES**

#### Example 1

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#### Production of phospholipase by cultivation of Hyphozyma

The strain *Hyphozyma sp.* CBS 648.91, was cultivated in a nutrient medium containing the following components:

Glucose 20 g/l Peptone 10 g/l MgSO<sub>4</sub>, 7 H<sub>2</sub>O 1 g/l Yeast Extract 10 g/l  $K_2HPO_4$  5 g/l

pH adjusted to 6.5 with NaOH

The strain was cultivated at 27-30 C for 3-4 days. The culture broth was subjected to liquid/solid separation by centrifugation. After centrifugation, a phospholipase activity of 1 unit/g culture broth was obtained (unit defined above). The supernatant was desalted and freeze-dried resulting in a crude powder preparation.

#### Example 2

#### Purification of phospholipase

Freeze dried phospholipase powder obtained according to Example 1 (300 units/g) was applied on a Butyl Toyopearl 650M column after adjusting the salt concentration to 3.5 M ammonium acetate. Bound phospholipase activity was eluted with H<sub>2</sub>O and separated from lipase activity which was also present in the crude powder preparation.

Fractions containing phospholipase activity were pooled, concentrated and dialyzed. The concentrated preparation was treated by anion exchange column chromatography using DEAE Toyopearl 650M. The adsorption condition was pH 7.5 (50 mM Tris-HCl) and elution was carried out by a linear gradient of 0-0.5M NaCl.

The last step was gel filtration column chromatography using HiLoad 26/60 Superdex 200pg. The condition was 50 mM Tris-HCl pH 7.5 including 0.5M NaCl. The resulting purified phospholipase was used in the following examples.

#### Example 3

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#### Characterization of phospholipase

The molecular weight (MW) of the phospholipase was found to be about 94 kDa on SDS PAGE and 87 kDa by gel filtration column chromatography. The polypeptide is believed to be glycosylated. The pl is around 5.6 on IEF PAGE.

The temperature profile was determined at pH 3.0 and 4.0 in a range of 40 to 70°C. The phospholipase was incubated for 10 minutes, and the activity was determined by the method described above. The temperature profile is presented in Fig. 1 as relative activity (taking the maximum activity as 100%). It is seen from this figure that both at pH 3 and 4, the phospholipase has high activity (more than 50% of optimum) at temperatures of 40 to 60°C with a temperature optimum around 50°C.

The pH profile was determined at 40°C using glycine-HCl buffer at pH 2, 2,5 and 3, and citrate buffer at pH 3, 4, 5 and 6. The results are presented in Fig. 2 as relative activity (taking the maximum activity as 100%). Due to a change of buffer system (glycine-HCl, citrate), the figure is made up of two curves, one representing the interval of pH 2.0 to 3.0 and the other representing the interval of pH 3.0 to 6.0. From the figure it appears that the phospholipase is active at pH values of 2 to 5, and the pH optimum is around 3.

The thermostability was determined by incubating in 0.1 M phosphate buffer (pH 7) for 10 minutes at temperatures of 40-80°C and determining the residual activity after the incubation. The results were 100% at 40°C, 95% at 50°C, 82% at 60°C, 55% at 70°C and 9% at 80°C. These results are also shown in Fig. 3.

#### 30 Example 4

#### Hydrolysis of phospholipid

A substrate solution was prepared by dissolving 2% of crude soy bean lecithin (phosphatidyl choline) in water. An enzyme solution was prepared by 50 times dilution of the purified enzyme from Example 2. 0.5 ml of the substrate solution, 0.25 ml of 0.4 M citrate buffer (pH 4) and 0.05 ml of 0.1 N CaCl<sub>2</sub> were mixed and incubated at 60°C.

0.1 ml of the enzyme solution was added and incubated for 1 hour at 60°C. The reaction was terminated by adding 0.1 ml of 1 N HCI. The mixture after the reaction was analyzed by TLC-latroscan as described above in the assay for reaction pattern.

The results showed that fatty acid was formed and that no lecithin remained after the reaction. A solid precipitate was observed at the bottom of the reaction vessel. This was believed to be a mixture of phospholipid and fatty acid.

#### Example 5

#### Hydrolysis of lyso-phospholipid

Lyso-phosphatidylcholine (LPC) was treated for 10 minutes at 40°C, other conditions being the same as described in Example 4. The chromatogram showed that about two thirds of the LPC was hydrolyzed, and that fatty acid was formed together with a small amount of phosphatidylcholine.

#### Example 6

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#### Enzymatic degumming of edible oil

Vegetable oil was degummed by treating it with phospholipase from *Hyphozyma* as follows. The enzyme dosage, the reaction pH and temperature were varied, and the resulting content of phospholipid was measured.

The equipment consisted of a 1 l jacketed steel reactor fitted with a steel lid, a propeller (600 rpm), baffles, a temperature sensor, an inlet tube at the top, a reflux condenser (4 °C) at the top, and an outlet tube at the bottom. The reactor jacket was connected to a thermostat bath. The outlet tube was connected via silicone tubing to an in-line mixer head equipped with a high shear screen (8500 rpm, flow ca. 1.1 l/minute). The mixer head was fitted with a cooling coil (5-10 °C) and an outlet tube, which was connected to the inlet tube of the reactor via silicone tubing. A temperature sensor was inserted in the silicone tubing just after the mixer head. The only connection from the reactor/mixer head system to the atmosphere was through the reflux condenser.

In each experiment, 0.6 I (ca. 560 g) of water-degummed rape seed oil with a P content of 186-252 ppm was loaded into the reactor with the thermostat and lab mixer running and pre-treated for 30 minutes with 0.6 g (2.86 mmol) of citric acid monohydrate in 27 g of water (added water vs. oil equals 4.8% w/w; [citric acid] in water phase = 106 mM, in water/oil emulsion = 4.6 mM) at time= 0. After the pre-treatment, the pH was adjusted by adding a NaOH solution followed by the enzyme solution. The mixture was then incubated for 6 hours, and samples for P-analysis and pH determination were drawn at intervals throughout the experiment.

The determination of phosphorous content in the oil was done according to procedure 2.421 in "Standard Methods for the Analysis of Oils, Fats, and Derivatives, 7.th ed. (1987)" after separating the emulsion by heating and centrifugation.

The initial performance was calculated from the initial rate of phosphorus 5 removal from the oil, taking the optimum as 100 %.

#### Degumming at various pH

The oil was treated at 40°C with an enzyme dosage of 1.3 mg/kg oil (as pure enzyme protein). The results at various pH were as follows:

pН	Initial performance	P content after 6 hours
	(relative to optimum)	
3.0	40	74 ppm
3.7	90	<10 ppm
4.4	100	<10 ppm
4.8	80	<10 ppm

#### Degumming at varioustemperatures

The oil was treated at pH 4.5 with an enzyme dosage of 1.3 mg/kg oil (as pure enzyme protein). The results at various temperatures were as follows:

Temperature	Initial performance	P content after 6 hours
	(relative to optimum)	
35°C	90	<10 ppm
40°C	100	<10 ppm

#### Degumming with various enzyme dosages

The oil was treated at pH 4.5, 40°C. The results at various enzyme dosages (given as pure enzyme protein) were as follows:

Enzyme dosage	Initial performance	P content after 6 hours
	(relative to optimum)	
0.65 mg/kg oil	70	<10 ppm
1.3	100	<10 ppm
2.6	100	<10 ppm

The results show good degumming performance at pH 3.5-5, 35-40°C. Good degumming to a phosphorus content below 10 ppm was obtained in 6 hours with a dosage of 1.3 mg/kg oil, and in 3 hours at a dosage of 2.6 mg/kg.

Measurement of the free fatty acids generated during degumming showed a low level of free fatty acids, corresponding very well to the amount of phospholipid in the substrate oil.

For reference, similar experiments were done with prior-art phospholipase from porcine pancreas. It was found that degumming to below 10 ppm of phosphorus could be obtained at 60°C, pH 5.5, but the performance of the prior-art enzyme dropped sharply at lower pH, and satisfactory degumming could not be achieved at pH lower than 5.5.

#### Example 7

#### Partial determination of the DNA sequence encoding the phospholipase

DNA encoding the phospholipase of *Hyphozyma* was isolated by two different methods. The 5' end of the gene was isolated by cloning. A genomic library of *Hyphozyma* DNA partially digested with Sau3A was screened at high stringency using standard methods (Sambrook et al., (1989), Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Lab.; Cold Spring Harbor, NY) with a probe specific to the phospholipase sequence. This probe was amplified from total *Hyphozyma* DNA with degenerate primers designed using the previously determined partial peptide sequences with SEQ ID NO: 1 and 5. Standard PCR conditions were used for amplification (Saiki et al., Science, 239, 487-491, 1988), including 0.5mM MgCl<sub>2</sub>, a 45°C annealing temperature, and primers PLMStr1 (SEQ ID NO: 12) and PLMStr6 (SEQ ID NO: 13). The clone pMStr16 hybridized to the probe, and therefore was isolated and a portion of the insert was sequenced.

An additional internal portion of the phospholipase-encoding gene was isolated using PCR with *Hyphozyma* DNA and the primers PLHaW2 (SEQ ID NO: 14) and PLMStr7 (SEQ ID NO: 15). PLHaW2 was designed using the sequence determined from pMStr16, and PLMStr7 was designed from the sequence of the partial peptide with SEQ ID NO: 8. Standard conditions were used for the PCR reactions, with 1.5 mM MgCl<sub>2</sub>, and a 46°C annealing temperature. The resulting amplified fragment was isolated and sequenced.

#### SEQUENCE LISTING

- (1) GENERAL INFORMATION:
  - (i) APPLICANT:
    - (A) NAME: Novo Nordisk A/S
    - (B) STREET: Novo Alle
    - (C) CITY: Bagsvaerd
    - (E) COUNTRY: Denmark
    - (F) POSTAL CODE (ZIP): DK-2880
    - (G) TELEPHONE: +45-4444-8888
    - (I) TELEX: +45-4449-3256
  - (ii) TITLE OF INVENTION: Novel Phospholipase, Production and Use Thereof
  - (iii) NUMBER OF SEQUENCES: 15
    - (iv) COMPUTER READABLE FORM:
      - (A) MEDIUM TYPE: Floppy disk
      - (B) COMPUTER: IBM PC compatible
      - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
      - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
- (2) INFORMATION FOR SEQ ID NO: 1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 16 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: N-terminal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

Ala Ser Pro Ser Gly Ser Tyr Ala Pro Ala Asn Met Pro Cys Xaa Gln

1 5 10 15

- (2) INFORMATION FOR SEQ ID NO: 2:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 7 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Asp Trp Ala Lys Trp Leu Ser

1

- (2) INFORMATION FOR SEQ ID NO: 3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 28 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

Asp Gly Arg Xaa Glu Thr Ala Asn Gln Arg Gly Thr Gly Gly Leu Leu

1 5 10 15

16

Gln Leu Ala Glu Tyr Ile Ala Gly Leu Ser Gly Gly
20 25

- (2) INFORMATION FOR SEQ ID NO: 4:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 36 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Asp Leu Glu Ser Asn Leu Ile Val Pro Glu Asp Gly Lys Val Ser Phe 1 5 10 15

Tyr Ala Ser Ile Leu Ala Ala Val Ala Gly Lys Arg Asn Glu Gly Tyr
20 25 30

Gln Thr Ser Leu

35

- (2) INFORMATION FOR SEQ ID NO: 5:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 41 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

Asp Glu Arg Glu Pro Gly Glu Leu Ile Ile Pro Arg Xaa Thr Thr Ile 5

10

15

Trp Glu Phe Asn Pro Tyr Glu Phe Gly Ser Trp Asn Pro Xaa Val Ser 20 25 30

Ala Phe Ile Pro Ile Glu Ile Leu Gly 35 40

- (2) INFORMATION FOR SEQ ID NO: 6:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
      - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) -ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

Asp Val Ser Leu Val Pro Asn Pro Phe Tyr Gly Tyr Val Gly Glu 5 10 15 1

- (2) INFORMATION FOR SEQ ID NO: 7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal

- (vi) ORIGINAL SOURCE:
  - (A) ORGANISM: Hyphozyma sp.
  - (B) STRAIN: CBS 648.91
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

Asp Val Thr Asn Trp Pro Xaa Ala Ser Ala Leu Tyr Gln Thr Ser Leu 1 5 10 15

Arg Ala Gln Tyr Pro Thr Tyr Ser Gln Tyr Ala Phe Pro Val 20 25 30

- (2) INFORMATION FOR SEQ ID NO: 8:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 31 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: internal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.
    - (B) STRAIN: CBS 648.91
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

Asp Thr Ser Phe Xaa Gly Thr Lys Thr Pro Ile Ile Val Tyr Met Pro 1 5 10 15

Ser Tyr Pro Tyr Ala Ala Phe Ala Asp Thr Ser Thr Phe Lys Leu 20 25 30

- (2) INFORMATION FOR SEQ ID NO: 9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 1870 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single

(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: DNA (genomic)

(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(v) FRAGMENT TYPE: N-terminal	
(vi) ORIGINAL SOURCE:	
(A) ORGANISM: Hyphozyma sp.	
(B) STRAIN: CBS 648.91	
(ix) FEATURE:	
(A) NAME/KEY: CDS	
(B) LOCATION:2141869	
(ix) FEATURE:	
(A) NAME/KEY: mat_peptide	
(B) LOCATION: 4421869	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:	
GGCGAGTGCA CAAGGCCGCG GACCAAATGT CCCTGAGTGC GTGTGTTTGT GTGTGACATA	60
GCCAGCAGAA TGCAGCTTAC TCTTCTTCCA TTGTGAGACG TTATATACCC ACACACATCT 1	20
CGCCGTCCCG TCAGACCCTT CTGCATCCGT CCGTACGAAC CTGCTCTCTT CCATTTACCT 1	80
CGACACTGTA TCGAGTGCAC GCTTCGAGGC ATC ATG AAG CTG CCG CTC CTC TCT 2	34
Met Lys Leu Pro Leu Leu Ser	
-76 -75 -70	
ACG CTG CTC AGC CTC GCG CTG ACC GCC TCG ACC GTC GTC CGT GCC TAT 2	82
Thr Leu Leu Ser Leu Ala Leu Thr Ala Ser Thr Val Val Arg Ala Tyr	
-65 -60 -55	
CCC TCC ATC CCG GCG CAG CTC ACC GAA GAC GAG ATC ACC CGC ATC AGC 3	30
Pro Ser Ile Pro Ala Gln Leu Thr Glu Asp Glu Ile Thr Arg Ile Ser	
-50 -45 -40	

CAG	CTC	TCC	CAG	GAG	GAC	AAG	GTC	AAG	TTT	GCC	GAA	CGC	ATC	CTA	GAG	378
Gln	Leu	Ser	Gln	Glu	Asp	Lys	Val	Lys	Phe	Ala	Glu	Arg	Ile	Leu	Glu	
		-35					-30					-25				
										•						
ATT	CGC	ACC	GCC	TAC	GAG	TAT	GAG	AAG	CAG	CAG	CTA	GCC	CGT	CAA	CAT	426
Ile	Arg	Thr	Ala	Tyr	Glu	Tyr	Glu	Lys	Gln	Gln	Leu	Ala	Arg	Gln	His	
	-20					-15					-10					
a.a.a	<b>a</b>			~~~	222	<b></b>			~~~							
			CGA													474
	ьеп	GIU	Arg	arg		ser	Pro	ser		ser	Tyr	Ala	Pro		Asn	
-5					1				5					10		
ATG	CCC	TGC	CCC	CAG	CGA	ACG	TCC	CAG	CAG	GGT	CCC	GGC	TTC	ATC	CGA	522
Met	Pro	Cys	Pro	Gln	Arg	Thr	Ser	Gln	Gln	Gly	Pro	Gly	Phe	Ile	Arg	
			15					20					25			
CCC	GCC	AAG	ACC	AAG	CAG	CTC	TCA	ATC	TCG	GAA	GCC	GAC	TAT	GTC	TCG	570
Pro	Ala	Lys	Thr	Lys	Gln	Leu	Ser	Ile	Ser	Glu	Ala	Asp	Tyr	Val	Ser	
		30					35					40				
CGC	CGC	CGC	ACC	AAC	ACC	CAG	GCC	GAC	TGG	GCC	AAG	TGG	CTC	TCG	GAC	618
Arg	Arg	Arg	Thr	Asn	Thr	Gln	Ala	Asp	Trp	Ala	Lys	Trp	Leu	Ser	Asp	
	45					50					55					
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TCG	TCG	ACC	GAC	CGC	GTG	CCT	CGT	CTG	GGC	TTT	GCG	CTC	AGC	GGC	GGT	714
Ser	Ser	Thr	Asp	Arg	Val	Pro	Arg	Leu	Gly	Phe	Ala	Leu	Ser	Gly	Gly	
				80					85					90		
GGA	CTG	CGT	GCC	ATG	CTC	GTT	GGT	TCG	GGC	ACG	CTC	CAG	GGC	TTT	GAC	762
Gly	Leu	Arg	Ala	Met	Leu	Val	Gly	Ser	Gly	Thr	Leu	Gln	Gly	Phe	Asp	
			95					100					105			

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GGC	CGC	AAC	GAG	ACC	GCC	AAC	CAG	CGT	GGC	ACC	GGT	GGA	CTG	CTC	CAG	810
Gly	Arg	Asn	Glu	Thr	Ala	Asn	Gln	Arg	Gly	Thr	Gly	Gly	Leu	Leu	Gln	
		110					115					120				
CTT	GCC	GAG	TAC	ATT	GCC	GGC	CTG	TCC	GGC	GGC	TCG	TGG	GCG	ACC	GCC	858
Leu	Ala	Glu	Tyr	Ile	Ala	Gly	Leu	Ser	Gly	Gly	Ser	Trp	Ala	Thr	Ala	
	125					130					135					
AGT	CTC	ACC	ATG	AAC	AAC	TGG	GCC	ACC	ACC	CAG	TCG	CTC	AAG	GAC	AAC	906
Ser	Leu	Thr	Met	Asn	Asn	Trp	Ala	Thr	Thr	Gln	Ser	Leu	Lys	Asp	Asn	
140					145					150					155	
ATC	TGG	GAT	CTC	GAG	TCC	AAC	CTC	ATC	GTC	CCC	GAG	GAC	GGC	AAG	GTC	954
Ile	Trp	Asp	Leu	Glu	Ser	Asn	Leu	Ile	Val	Pro	Glu	Asp	$\operatorname{Gl}_{\mathbf{Y}}$	Lys	Val	
				160					165					170		
TCG	TTT	TAC	GCC	TCG	ATC	CTG	GCC	GCC	GTC	GCG	GGC	AAG	AGG	AAC	GAA	1002
Ser	Phe	Tyr	Ala	Ser	Ile	Leu	Ala	Ala	Val	Ala	Gly	Lys	Arg	Asn	Glu	
			175					180					185			
GGT	TAC	CAG	ACC	AGT	CTC	ACC	GAC	TAC	TTT	GGC	CTC	TCG	ATC	GCC	GAC	1050
Gly	Tyr	Gln	Thr	Ser	Leu	Thr	Asp	Tyr	Phe	Gly	Leu	Ser	Ile	Ala	Asp	
		190					195					200				
AAG	ATT	CTC	AAC	GGC	TCC	ATG	TAC	GGC	AAC	AAG	TTC	AGC	GTC	GAG	TGG-	1098
Lys	Ile	Leu	Asn	Gly	Ser	Met	Tyr	Gly	Asn	Lys	Phe	Ser	Val	Glu	Trp	
	205					210					215					
AGC	GAC	GTC	AAG	AAT	ACG	TCC	AAG	TTC	ACC	GAT	GCC	TCC	ATG	CCG	TTC	1146
Ser	Asp	Val	Lys	Asn	Thr	Ser	Lys	Phe	Thr	Asp	Ala	Ser	Met	Pro	Phe	•
220					225					230					235	
CCC	ATC	ATT	ATT	GCC	GAC	GAG	CGC	GAG	CCC	GGC	GAG	CTC	ATC	ATC	CCG	1194
Pro	Ile	Ile	Ile	Ala	Asp	Glu	Arg	Glu	Pro	Gly	Glu	Leu	Ile	Ile	Pro	
				240					245	_				250		

CGC	AAC	ACC	ACC	ATC	TGG	GAG	TTC	AAC	CCG	TAC	GAG	TTC	GGT	TCT	TGG	1242
Arg	Asn	Thr	Thr	Ile	Trp	Glu	Phe	Asn	Pro	Tyr	Glu	Phe	Gly	Ser	Trp	
			255					260					265			
AAC	CCC	AAT	GTT	TCG	GCT	TTC	ATC	CCC	ATC	GAG	ATC	CTC	GGC	TCG	AGT	1290
Asn	Pro	Asn	Val	Ser	Ala	Phe	Ile	Pro	Ile	Glu	Ile	Leu	Gly	Ser	Ser	
		270					275					280				
CTG	GAC	AAC	GGC	ACC	AGC	GTC	CTG	CCC	GAC	GGC	GTC	TGT	GTC	GGC	GGA	1338
Leu	Asp	Asn	Gly	Thr	Ser	Val	Leu	Pro	Asp	Gly	Val	Cys	Val	Gly	Gly	
	285					290					295					
TAC	GAG	ACC	GTT	GCC	TGG	GTG	ACT	GGC	ACC	TCG	GCG	ACT	CTG	TTC	TCT	1386
Tyr	Glu	Thr	Val	Ala	Trp	Val	Thr	Gly	Thr	Ser	Ala	Thr	Leu	Phe	Ser	
300					305					310					315	
GGT	CTG	TAC	CTC	GAA	CTT	ATC	TCG	ACC	TCG	AGC	AAC	AAC	ATC	ATC	GTC	1434
Gly	Leu	Tyr	Leu	Glu	Leu	Ile	Ser	Thr	Ser	Ser	Asn	Asn	Ile	Ile	Val	
				320					325					330		
GAT	GCG	CTC	AAG	GAG	ATT	GCC	CAG	GCG	GTA	TCA	AAC	GAG	CAG	AAC	GAT	1482
Asp	Ala	Leụ	Lys	Glu	Ile	Ala	Gln	Ala	Val	Ser	Asn	Glu	Gln	Asn	Asp	
			335					340					345			
														GGC	*	1530
Val	Ser		Val	Pro	Asn	Pro	Phe	Tyr	Gly	Tyr	Val	Gly	Glu	Gly	Asp	
		350					355					360				
														GGT		1578
Val		Val	Ser	Asp	Leu	Arg	Asn	Ile	Thr	Leu	Val	Asp	Gly	Gly	Leu	
	365					370					375					
														CGC		1626
	Asn	Glu	Asn	Val		Leu	Trp	Pro	Leu		Glu	Pro	Ala	Arg	Asp	
380					385					390					395	

CTG	GAC	GTG	ATC	ATC	GCC	ATT	GAC	AGC	TCG	GCG	GAC	GTG	ACC	AAC	TGG	1674
Leu	Asp	Val	Ile	Ile	Ala	Ile	Asp	Ser	Ser	Ala	Asp	Val	Thr	Asn	Trp	
				400					405					410		
CCG	AAC	GCG	TCG	GCG	CTG	TAC	CAG	ACG	TCG	CTG	CGT	GCT	CAG	TAC	CCG	1722
Pro	Asn	Ala	Ser	Ala	Leu	Tyr	Gln	Thr	Ser	Leu	Arg	Ala	Gln	Tyr	Pro	
			415					420					425			
3.00	m > m		G 2 G		~~~		~~~	~~~								
		AGC														1770
IIIL	ıyı	Ser	GIII	ıyr	Ата	Pne	435	vai	мес	Pro	Asp		Asn	Thr	Val	
		430					433					440				
GTC	AAC	CGC	GGC	CTC	AAC	ACG	CGC	CCC	GTG	TTC	TAC	GGC	TGC	ААТ	GCG	1818
		Arg														1010
	445		•			450	J				455	•	-			
ACC	GTC	AAC	GTC	ACC	AAC	GCG	GAT	ACG	TCG	TTC	AAC	GGC	ACC	AAG	ACG	1866
Thr	Val	Asn	Val	Thr	Asn	Ala	Asp	Thr	Ser	Phe	Asn	Gly	Thr	Lys	Thr	
460					465					470					475	
CCA	A															1870
Pro			-													
(2)	TNE	ר זאנטר	PT (N)	EOB	CEO	TD N	. Ot	10.								
(2)		ORMAT (i) S														
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		(1	) T(	OPOLO	OGY:	line	ear									
	(ii)	) MOI	LECUI	LE T	YPE:	prot	cein									
	(xi	) SE	QUEN	CE DI	ESCR	IPTIC	ON: S	SEQ :	ID NO	): 10	):					
		Leu	Pro	Leu	Leu	Ser	Thr	Leu	Leu	Ser	Leu	Ala	Leu	Thr	Ala	
-76	-75					-70					-65					
0	ml	** 7		40		<b></b>		•		_		<b>~</b> 3	_		-3	
	Tnr	Val	val	Arg		Tyr	Pro	ser	Пе		Ala	GIn	Leu	Thr		
-60					-55					-50					-45	

Asp	Glu	Ile	Thr		Ile	Ser	Gln	Leu		Gln	Glu	Asp	Lys	Val	Lys
				-40					-35					-30	
Phe	Ala	Glu	Arg	Ile	Leu	Glu	Ile	Arg	Thr	Ala	Tyr	Glu	Tyr	Glu	Lys
			-25					-20					-15		
Gln	Gln	Leu	Ala	Arg	Gln	His	Ala	Leu	Glu	Arg	Arg	Ala	Ser	Pro	Ser
		-10					-5					1			
Gly	Ser	Tyr	Ala	Pro	Ala	Asn	Met	Pro	Cys	Pro	Gln	Arg	Thr	Ser	Gln
5		_			10				_	15					20
Gln	Glv	Pro	Gly	Phe	Ile	Ara	Pro	Ala	īvs	Thr	Lvs	Gln	Len	Ser	Tle
	1		1	25		3			30		-1-			35	
Con	Cl.	71-	7 ~~		val.	C0~	7~~	y.c.	λνα	Wha	7 an	ሞኮሎ	<b>61</b> 2	71-	) an
ser	GIU	AIA	Asp 40	TÄT	Val	ser	Arg	45	Arg	Inr	ASII	IIII	50	АТА	Asp
										_	_				
Trp	Ala	Lys 55	Trp	Leu	Ser	Asp	Ser 60	Ala	Lys	Leu	Asn	Ser 65	Ser	Leu	Pro
											•				
Gly	Gly 70	Ala	Ser	Asn	Tyr	Thr 75	Ser	Ser	Thr	Asp	Arg 80	Val	Pro	Arg	Leu
	, 0					, ,									
_	Phe	Ala	Leu	Ser	_	Gly	Gly	Leu	Arg		Met	Leu	Val	Gly	
85					90					95					100
Gly	Thr	Leu	Gln	_	Phe	Asp	Gly	Arg		Glu	Thr	Ala	Asn		Arg
				105					110					115	
Gly	Thr	Gly	Gly	Leu	Leu	Gln	Leu	Ala	Glu	Tyr	Ile	Ala	Gly	Leu	Ser
			120					125					130		
Gly	Gly	Ser	Trp	Ala	Thr	Ala		Leu	Thr	Met	Asn	Asn	Trp	Ala	Thr
		135					140					145			

Thr	Gln 150	Ser	Leu	Lys	Asp	Asn 155	Ile	Trp	Asp	Leu	Glu 160	Ser	Asn	Leu	Ile
Val	Pro	Glu	Asp	Gly	Lys 170	Val	Ser	Phe	Tyr	Ala 175	Ser	Ile	Leu	Ala	Ala 180
Val	Ala	Gly	Lys	Arg 185	Asn	Glu	Gly	Tyr	Gln 190	Thr	Ser	Leu	Thr	Asp 195	Tyr
Phe	Gly	Leu	Ser 200	Ile	Ala	Asp	Lys	Ile 205	Leu	Asn	Gly	Ser	Met 210	Tyr	Gly
Asn	Lys	Phe 215	Ser	Val	Glu	Trp	Ser 220	Asp	Val	Lys	Asn	Thr 225	Ser	Lys	Phe
Thr	Asp 230	Ala	Ser	Met	Pro	Phe 235	Pro	Ile	Ile	Ile	Ala 240	Asp	Glù	Arg	Glu
Pro 245	Gly	Glu	Leu	Ile	Ile 250	Pro	Arg	Asn	Thr	Thr 255	Ile	Trp	Glu	Phe	Asn 260
Pro	Tyr	Glu	Phe	Gly 265	Ser	Trp	Asn	Pro	Asn 270	Val	Ser	Ala	Phe	Ile 275	Pro
Ile	Glu	Ile	Leu 280	Gly	Ser	Ser	Leu	Asp 285	Asn	Gly	Thr	Ser	Val 290	Leu	Pro
Asp	Gly	Val 295	Cys	Val	Gly	Gly	Tyr 300	Glu	Thr	Val	Ala	Trp 305	Val	Thr	Gly
Thr	Ser 310	Ala	Thr	Leu	Phe	Ser 315	Gly	Leu	Tyr	Leu	Glu 320	Leu	Ile	Ser	Thr
Ser 325	Ser	Asn	Asn	Ile	Ile 330	Val	Asp	Ala	Leu	Lys 335	Glu	Ile	Ala	Gln	Ala

Val	Ser	Asn	Glu	Gln	Asn	Asp	Val	Ser	Leu	Val	Pro	Asn	Pro	Phe	Tyr
				345					350					355	

- Gly Tyr Val Gly Glu Gly Asp Val Gln Val Ser Asp Leu Arg Asn Ile 360 365 370
- Thr Leu Val Asp Gly Gly Leu Asp Asn Glu Asn Val Pro Leu Trp Pro 375 380 385
- Leu Val Glu Pro Ala Arg Asp Leu Asp Val Ile Ile Ala Ile Asp Ser 390 395 400
- Ser Ala Asp Val Thr Asn Trp Pro Asn Ala Ser Ala Leu Tyr Gln Thr 405 410 415 420
- Ser Leu Arg Ala Gln Tyr Pro Thr Tyr Ser Gln Tyr Ala Phe Pro Val 425 430 435
- Met Pro Asp Thr Asn Thr Val Val Asn Arg Gly Leu Asn Thr Arg Pro 440 445 450
- Val Phe Tyr Gly Cys Asn Ala Thr Val Asn Val Thr Asn Ala Asp Thr
  455 460 465

Ser Phe Asn Gly Thr Lys Thr Pro 470 475

- (2) INFORMATION FOR SEQ ID NO: 11:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 573 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: peptide
  - (v) FRAGMENT TYPE: N-terminal
  - (vi) ORIGINAL SOURCE:
    - (A) ORGANISM: Hyphozyma sp.

70

	(B)	ST	RAIN	CBS	5 648	3.91									
(ix)	FEAT	TURE:	:												
	(A)	(A) NAME/KEY: Protein													
	(B)	LO	CATIO	ON:1	497	7									
(xi)	SEQU	JENCI	E DES	SCRI	PTION	1: SI	EQ II	OM C	: 11	:					
Met	Lys	Leu	Pro	Leu	Leu	Ser	Thr	Leu	Leu	Ser	Leu	Ala	Leu	Thr	Ala
	-75					-70					-65				
	Thr	Val	Val	Arg	Ala	Tyr	Pro	Ser	Ile	Pro	Ala	Gln	Leu	Thr	Glı
-60					-55					-50					-4!
Asp	Glu	Ile	Thr		Ile	Ser	Gln	Leu	Ser	Gln	Glu	Asp	Lys	Val	Ly
				-40					-35					-30	
				_		_									
Phe	Ala	Glu		Ile	Leu	Glu	Ile		Thr	Ala	Tyr	Glu	Tyr	Glu	Lys
			-25					-20					-15		
		_		_											
Gin	Gln		Ala	Arg	GIn	His		Leu	Glu	Arg	Arg	Ala	Ser	Pro	Sei
		-10					-5					1			
Clv	Cor	T1 170	ח ה	Dro	ת ות	7 an	Mot	Dro	C***	Dwo	<b>71</b> -	λ	mb	0	~1-
	Ser	ıyı	Ala	PIO		ASII	Mec	PIO	cys		GIII	Arg	1111	Ser	
5					10					15					20
Gln	Gly	Dro	Glv	Dhe	Tla	λνα	Pro	כות	Larc	Thr	Larc	Cln.	Lou	Cor	Tla
GIII	Gry	FIO	Gly		116	лц	FIO	AIG	30	1111	пуъ	GIII	neu		116
				25					30					35	
Ser	Glu	Δla	Asn	ጥህጕ	Val	Ser	Δrα	Δτα	Δνα	Thr	Λcn	Thr	Gln	בות	λcr
202			40	-1-		001		45	*** 9		71011	****	50	niu	TOF
													50		
Tro	Ala	Lvs	Tro	Leu	Ser	Asp	Ser	Ala	Lvs	Len	Asn	Ser	Ser	I,en	Pro
		55					60		10			65	JUL	2,04	
												<b>.</b> .			

Gly Gly Ala Ser Asn Tyr Thr Ser Ser Thr Asp Arg Val Pro Arg Leu

80

75

Gl <sub>y</sub> 85	Phe	Ala	Leu	Ser	Gly 90	Gly	Gly	Leu	Arg	Ala 95	Met	Leu	Val	Gly	Ser
Gly	Thr	Leu	Gln	Gly 105	Phe	Asp	Gly	Arg	Asn 110	Glu	Thr	Ala	Asn	Gln 115	Arg
Gly	Thr	Gly	Gly 120	Leu	Leu	Gln	Leu	Ala 125	Glu	Tyr	Ile	Ala	Gly 130	Leu	Ser
Gly	gly	Ser 135	Trp	Ala	Thr	Ala	Ser 140	Leu	Thr	Met	Asn	Asn 145	Trp	Ala	Thr
Thi	Gln 150	Ser	Leu	Lys	Asp	Asn 155	Ile	Trp	Asp	Leu	Glu 160	Ser	Asn	Leu	Ile
Va]	Pro	Glu	Asp	Gly	Lys 170	Val	Ser	Phe	Tyr	Ala 175	Ser	Ile	Leu	Ala	Ala 180
Va]	. Ala	Gly	Lys	Arg 185	Asn	Glu	Gly	Tyr	Gln 190	Thr	Ser	Leu	Thr	Asp 195	Tyr
Phe	e Gly	Leu	Ser 200	Ile	Ala	Asp	Lys	Ile 205	Leu	Asn	Gly	Ser	Met 210	туг	Gly
Ası	ı Lys	Phe 215	Ser	Val	Glu	Trp	Ser 220	Asp	Val	Lys	Asn	Thr 225	Ser	Lys	Phe
Thi	230	Ala	Ser	Met	Pro	Phe 235	Pro	Ile	Ile	Ile	Ala 240	Asp	Glu	Arg	Glu
Pro 24!	o Gly	Glu	Leu	Ile	Ile 250	Pro	Arg	Asn	Thr	Thr 255	Ile	Trp	Glu	Phe	Asn 260
Pro	o Tyr	Glu	Phe	Gly 265	Ser	Trp	Asn	Pro	Asn 270	·Val	Ser	Ala	Phe	Ile 275	Pro

WO 98/18912

Ile	Glu	Ile	Leu	Gly	Ser	Ser	Leu	Asp	Asn	Gly	Thr	Ser	Val	Leu	Pro
			280					285					290		
Asp	Glv	Val	Cvs	Val	Glv	Glv	Tvr	Glu	Thr	Val	Ala	Tro	Val	Thr	Glv

Asp Gly Val Cys Val Gly Gly Tyr Glu Thr Val Ala Trp Val Thr Gly
295 300 305

Thr Ser Ala Thr Leu Phe Ser Gly Leu Tyr Leu Glu Leu Ile Ser Thr
310 315 320

Val Ser Asn Glu Gln Asn Asp Val Ser Leu Val Pro Asn Pro Phe Tyr 345 350 355

Gly Tyr Val Gly Glu Gly Asp Val Gln Val Ser Asp Leu Arg Asn Ile 360 365 370

Thr Leu Val Asp Gly Gly Leu Asp Asn Glu Asn Val Pro Leu Trp Pro 375 380 385

Leu Val Glu Pro Ala Arg Asp Leu Asp Val Ile Ile Ala Ile Asp Ser 390 395 400

Ser Ala Asp Val Thr Asn Trp Pro Asn Ala Ser Ala Leu Tyr Gln Thr 405 410 415 420

Ser Leu Arg Ala Gln Tyr Pro Thr Tyr Ser Gln Tyr Ala Phe Pro Val 425 430 435

Met Pro Asp Thr Asn Thr Val Val Asn Arg Gly Leu Asn Thr Arg Pro 440 445 450

Val Phe Tyr Gly Cys Asn Ala Thr Val Asn Val Thr Asn Ala Asp Thr
455 460 465

30

Ser Phe Asn Gly Thr Lys Thr Pro Ile Ile Val Tyr Met Pro Ser Tyr 470 475 480

Pro Tyr Ala Ala Phe Ala Asp Thr Ser Thr Phe Lys Leu 485 490 495

- (2) INFORMATION FOR SEQ ID NO: 12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid
  - (ix) FEATURE:
    - (A) NAME/KEY: modified base
    - (B) LOCATION:3..18
    - (D) OTHER INFORMATION:/mod\_base= OTHER /note= "deoxyinosine"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

GCNCCNGCNA AYATGCCNTG

20

- (2) INFORMATION FOR SEQ ID NO: 13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 20 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: other nucleic acid
  - (ix) FEATURE:
    - (A) NAME/KEY: modified\_base
    - (B) LOCATION:6
    - (D) OTHER INFORMATION:/mod\_base= OTHER
      /note= "deoxyinosine"
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GGCATGTAGA CGATGAT

17

TCGT	FANGGGT TRAAYTCCCA	20
(2)	INFORMATION FOR SEQ ID NO: 14:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
CCAT	TGCTCGT TGGTTCG	17
(2)	INFORMATION FOR SEQ ID NO: 15:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 17 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	

## INDICATIONS RELATING TO A DEPOSITED MICROORGANISM (PCT Rule 13bis)

A. The indications made below relate to the microorganism referred to in the description on page 6, lines 18-22.									
B. IDENTIFICATION OF DEPOSIT F	urther deposits are identified on an additional sheet								
Name of depositary institution									
Centraal Bureau voor Schimmelcultures (CBS)									
Address of depositary institution (including postal code and country)									
Oosterstraat 1, 3740 AG Barrn, Netherlands									
Date of deposit 12 November 1991	Accession Number CBS 648.91								
	C. ADDITIONAL INDICATIONS (leave blank if not applicable) This information is continued on an additional sheet								
withdrawn, a sample of the deposited microorganism is only to be provided to an independent expert nominated by the person requesting the sample (cf. Rule 28(4) EPC). As far as Australia is concerned, the expert option is likewise requested, reference being had to Regulation 3.25 of Australia Statutory Rules 1991 No 71. Also, for Canada we request that only an independent expert nominated by the Commissioner is authorized to have access to a sample of the microorganism deposited.  D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (if the indications are not for all designated States)									
E. SEPARATE FURNISHING OF INDICATIONS (leave blank if not applicable)									
The indications listed below will be submitted to the International Bureau later (specify the general nature of the indications e.g., "Accession Number of Deposit")									
For receiving Office use only	For International Bureau use only								
This sheet was received with the international application	This sheet was received by the International Bureau on:								
Authorized officer	Authorized officer								
Trom Gualania.									

FORM PCT/RO/134 (JULY 1992)

### **CLAIMS**

- 1. A phospholipase which:
  - a) is able to hydrolyze both fatty acyl groups in a phospholipid,
  - b) is derivable from a strain of *Hyphozyma*,
- 5 c) has a temperature optimum measured for 10 minutes at pH 3-4 of about 50°C, and
  - d) has a pH optimum measured at 40°C for 10 minutes of about pH 3.
  - 2. A phospholipase which:
    - a) is able to hydrolyze both fatty acyl groups in a phospholipid, and
- b) is a polypeptide comprising an N-terminal amino acid sequence which is the sequence shown in positions 1-497 of SEQ ID NO: 11, or is at least 50 % identical therewith.
  - 3. A phospholipase which:
    - a) is able to hydrolyze both fatty acyl groups in a phospholipid, and
- b) is a polypeptide containing amino acid sequences which are at least 50% identical with the amino acid sequences shown in SEQ ID NO: 1-8.
  - 4. The phospholipase of claim 2 or 3 wherein said identity of sequences is at least 60%, preferably at least 70%, more preferably at least 80%, and most preferably at least 90%.
- The phospholipase of any preceding claim which is derivable from *Hyphozyma* sp. strain CBS 648.91.
  - 6. The phospholipase of any preceding claim which is essentially free from lipase activity.
  - 7. A DNA sequence which encodes the phospholipase of claim 2.
- 25 8. The DNA sequence of the preceding claim which comprises the sequence shown in positions 457-1870 of SEQ ID NO: 9.

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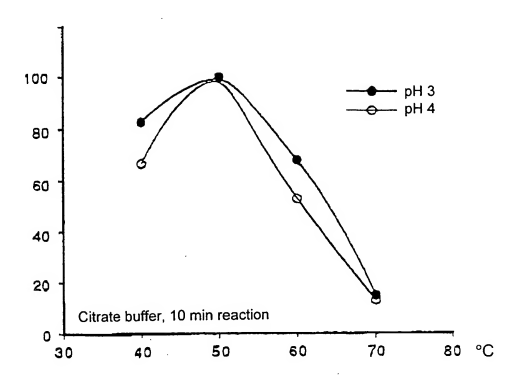
- 9. A method of producing a phospholipase, comprising cultivation of a phospholipase-producing strain of *Hyphozyma* in a suitable nutrient medium, followed by recovery of the phospholipase.
- 10. The method of the preceding claim wherein the strain is *Hyphozyma sp.* strain 5 CBS 648.91.
  - 11. The method of claim 9 or 10 wherein the recovery comprises separation to remove lipase activity.
  - 12. A method for producing a phospholipase, comprising:
    - a) isolating a DNA sequence encoding the phospholipase from a phospholipase-producing strain of *Hyphozyma*,
    - b) combining the DNA fragment with appropriate expression signal(s) in an appropriate vector,
    - c) transforming a suitable heterologous host organism with the vector,
    - d) cultivating the transformed host organism under conditions leading to expression of the phospholipase, and
    - e) recovering the phospholipase from the culture medium.
- 13. The method of the preceding claim, wherein the host organism is a eukaryotic cell, in particular a fungal cell, such as a yeast cell or a filamentous fungal cell, preferably a strain of Aspergillus, Fusarium, Trichoderma or Saccharomyces, most preferably A. niger, A. oryzae, F. graminearum, F. sambucinum, F. cerealis or S. cerevisiae.
  - 14. The method of claim 12 or 13 wherein the DNA sequence is isolated by a method comprising:
    - a) cloning, in suitable vectors, a cDNA library from the phospholipase-producing strain of *Hyphozyma*,
    - b) transforming suitable yeast host cells with said vectors,
    - c) cultivating the transformed yeast host cells under suitable conditions to express the phospholipase,
- d) screening for positive clones by determining the phospholipase activity expressed in step (c).

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- 15. The method of any of claims 12-14, wherein the *Hyphozyma* strain is *Hyphozyma sp.* strain CBS 648.91.
- 16. A process for hydrolyzing fatty acyl groups in a phospholipid or lysophospholipid, comprising treating the phospholipid or lysophospholipid with the 5 phospholipase of any of claims 1-6.
  - 17. The process of the preceding claim wherein the phospholipid or lysophospholipid comprises lecithin or lysolecithin.
  - 18. The process of claim 16 or 17 wherein the treatment is conducted at pH 1.5-5 (preferably 2-4) and 30-70°C.
- 10 19. The process of any of claims 16-18, which is a process for improving the filterability of an aqueous solution or slurry of carbohydrate origin which contains phospholipid.
  - 20. The process of the preceding claim wherein the solution or slurry contains a starch hydrolysate, particularly a wheat starch hydrolysate.
- 15 21. The process of any of claims 16-18 which is a process for making bread, comprising adding the phospholipase to the ingredients of a dough, kneading the dough and baking the dough to make the bread.
- 22. The process of any of claims 16-18 which is a process for reducing the content of phospholipid in an edible oil, comprising treating the oil with the phospholipase so as to hydrolyze a major part of the phospholipid, and separating an aqueous phase containing the hydrolyzed phospholipid from the oil.
  - 23. A process for removing phospholipid from an edible oil, comprising:
    - a) treating the oil at pH 1.5-3 with a dispersion of an aqueous solution of a phospholipase having the ability to hydrolyze the intact phospholipid at said pH, so as to hydrolyze a major part of the phospholipid, and
    - b) separating an aqueous phase containing the hydrolyzed phospholipid from the oil.

- 24. The method of the preceding claim wherein the oil is treated to remove mucilage prior to the treatment with the phospholipase.
- 25. The method of claim 23 or 24 wherein the oil prior to the treatment with the phospholipase contains the phospholipid in an amount corresponding to 50-250 ppm 5 as phosphorus.
  - 26. The method of any of claims 23-25 wherein the phospholipase is the phospholipase of any of claims 1-6.
- 27. The process of any of claims 23-26 wherein the treatment with phospholipase is done at 30-45°C for 1-12 hours at a phospholipase dosage of 0.1-10 mg/l in the 10 presence of 0.5-5% of water.

Relative activity (%)



Temperature profile

FIG. 1

Relative activity (%)

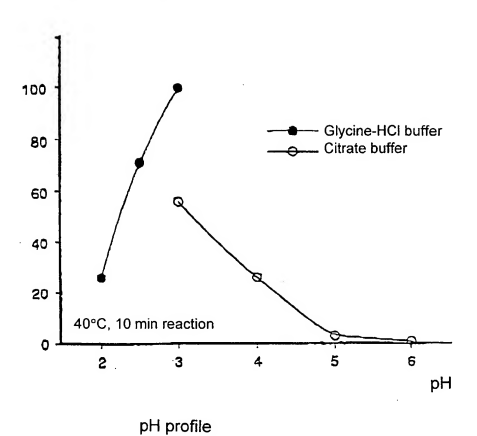


FIG. 2

# Remaining activity (%)

Thermostability

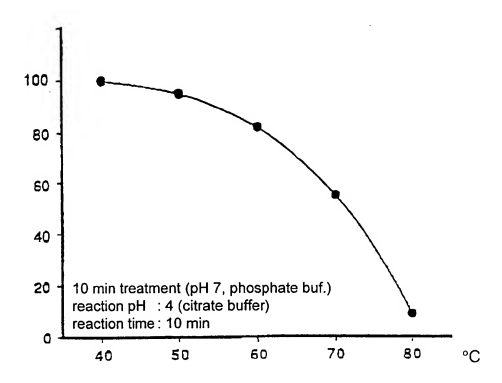


FIG. 3

09	[LEIRTAY	120   	180   
50	)EDKVKFAER]	110     o o   PPAKTKQLSIS REASGLSDN QASGPSDN SSAAKLSTN	170 1     o ooo oo   o o   FALSGGGLRAMLVGS(   VACSGGGYRAMLSGA(   VACSGGGYRAMLSGA(
04 —	DEITRISQLSC		160     o   o STDRVPRLGFP NSSNMPKIAVP NASDIPRIAVP
30	YPSIPAQLTEI NAWS LAWS LPNA	90 	150    - 
70	STLLSLALTASTVVRAYPSIP; SLLVSAAVLTSLTENVNAWS EWLLFSDAVFFAQG-TLAWS	70   0 0 0 0 	140    SDSAKLNSSI  INRATSNFSD
10	MKLPLLSTLLSLALTASTVVRAYPSIPAQLTEDEITRISQLSQEDKVKFAERILEIRTAY MKLQSLLVSAAVLTSLTENVNAWS	70 70 90 100 110 120	130 140 150 160 170 180
	Hyphozyma Saccharomyces Torulaspora Penicillium	Hyphozyma Saccharomyces Torulaspora Penicillium	Hyphozyma Saccharomyces Torulaspora Penicillium

FIG. 4a

240	0	MIN	TTESNSIW	MTEDDSIW	EAGSVW
230		ATTQSLKD-	TSVQAIVDN	TSVQDIVNN	ТТVDКLQТН
220	-   0   -	WATASLTMINIW	WLTSTLAWNNW	WLVGTLAWNNW	WLLGSIYINNF
210	0 00000 0	<b>AEYIAGLSGGS</b>	ATYLAGLSGGN	TTYLAGLSGGNV	ATYISGLSGGS
200	00000	3-TGGLLQL	3-LGGLLQG	3-LGGLLQS	SHLGGLLQS
190	0 00000 0 0 0 0 0 0	GFDGRNETANQRG-TGGLLQLAEYIAGLSGGSWATASLTMNNWATTQSLKDNIW	AMDNRTDGANEHG-LGGLLQGATYLAGLSGGNWLTSTLAWNNWTSVQAIVDNTTESNSIW	AMDNRTDGANEHG-LGGLLQSTTYLAGLSGGNWLVGTLAWNNWTSVQDIVNNMTEDDSIW	AFDSRTDNATATGHLGGLLQSATYISGLSGGSWLLGSIYINNFTTVDKLQTHEAGSVW
		Hyphozyma	Saccharomyces	Torulaspora	Penicillium

300	KILNGSM
290	o     DYFGLSIAD
0	o  o   o o agkrnegyǫtslti
280	o o ol
270	, o c KVSFYASILAAVA
260	EDG-KVS
250	o    o DLESNLIV-P-

DISNSIIN-P--GGFMIVTTIKRWDHISDAVEGKQDAGFNVSLTDIWGRALSYNFFPSLY DISHSILT-P--DGINIFKTGSRWDDISDDVQDKKDAGFNISLADVWGRALAYNFWPSLH QFGNSIIEGPDAGGIQLLDSAGYYKDLADAVDGKKKAGFDTTLTDIWGRALSYQMFNASN Saccharomyces

Penicillium

Torulaspora

Hyphozyma

R-GGVAYTWSTLRDVEVFQNGEMPFPISVADGRYPGTQIIDLNATVFEFNPFEMGSWDPT --GGLSYTWSSIADTPEFQDGDYPMPFVVADGRNPGELVIGSNSTVYEFNPWEFGTFDPT Saccharomyces Torulaspora Penicillium Hyphozyma

420	ISSNNII FOLPSFI FSLPSFI	480   0   00 DNENVPL DNQNIPL DGQNVPL	540
410   0   0	SGLILELLS NQFLLRLNS: NQFLLQINS: NQFLLQINT:	470 48C 	530
400 	AWVIGISALLE GFITATSSTLE GFIMGTSSSLE GFVIGTSSSLE	460    -EGDVQVSDLF  KNATSSIIESE  DNFSKSISESE  -EHSSPYAAQF	520
390	DGVCVGGIEIV KGQCIAGFDNT KGQCVAGYDNT ES-CIRGFDSP	450   oo PNPFYGYVG PNPFKEANFLÇ PNPFKDTSYIÇ	510
380	SLUNGISVLF NVTNG-KPVN KVSNG-EPVN KFEGGSLPSN	440   0000   NEQNDVSLV-PN NNSDDIAIYAPN SDEDDIAIYAPN KSQNDIASYDPN	500
370	VSAFIPIELLGSSLDNGISVLPDGVCVGGIEIVAWVIGISAILFSGLILELISISSNALL LNAFTDVKYLGTRVSNG-KPVNKGQCIAGFDNTGFITATSSTLFNQFLLRLNSTDLPSFI LNAFTDVKYLGTKVSNG-EPVNKGQCVAGYDNTGFIMGTSSSLFNQFLLQINSTSLPSFI IFGFVPLEYLGSKFEGGSLPSNES-CIRGFDSAGFVIGTSSSLFNQFLLQINTTSLPSFI	430 440 450 460 470 480	4 90
,	Hyphozyma Saccharomyces Torulaspora Penicillium	Hyphozyma Saccharomyces Torulaspora Penicillium	

FIG. 4c

VPLLQKERELDVIFALDNSADTDDYWPDGASLVNTYQRQFGSQGLN--LSFPYVPDVNTF VPLVQDERNVDVIFALDNSADTDYYWPDGASLVSTYERQFSSQGLN--MSFPYVPDKRTF HPLIQPERHVDVIFAVDSSADTDYFWPNGTSLVATYERSLNSSGIANGTAFPAVPDQNTF

Saccharomyces Torulaspora Penicillium

Hyphozyma

WPLVEPARDLDVIIAIDSSADVTN-WPNASALYQTSLRAQYPTYSQ--YAFPVMPDTNTV

0

0

0

00 00

600     TSTFKL QSTFKMSYSDSE TSTFKLSYTDDE ISTFQLSTDDAE	099	STINYCWNGTI SFINYCWNGTI	720	NAGNALVNYSNL -AGLPA L		Q
590   OC YAAFADTS HSYNGNQS	650	TPSECSOC TPECSTC	710	SSTHKKN		
	640		700			F H C
570    NADTSFNG:  DLEYIP	630	 DFLGCVGC/ SFMGCVAC/	069	 SAAAASASI YSAYNTES! DPAFYLADI		
550 560 570 570 570 570 570 570 570 570 570 57	620	 AATMGNFTDDS AATRGNLTDDS VATMANSTLDD	680	NDDYSSSASLSASAAASASASASASASA NSDFDPTAASSAYSAYNTESYSSSSATGS		VISAVFGLI LLTAIAGFL GLAMLILV-
550    o oo  o o VNRGLNTRPV; VNLGLNKKPT; VNLGLADKPS; INLGLSTRPS;	610	RLGMIKNGFEA RLKMIKNGFEA RDNIILNGYEV	670	DSRSVSGVGN: DDTPVSGLDN	730	NTNTFIGVLSVISAVFGLI TPTSFTSILTLLTAIAGFL PTMLSTVVAAGLAMLILV-
Hyphozyma Saccharomyces Torulaspora Penicillium		Hyphozyma Saccharomyces Torulaspora Penicillium		Hyphozyma Saccharomyces Torulaspora Penicillium		Hyphozyma Saccharomyces Torulaspora Penicillium

International application No. PCT/DK 97/00490 A. CLASSIFICATION OF SUBJECT MATTER IPC6: C12N 9/18, C11B 3/00 According to International Patent Classification (IPC) or to both national classification and IPC

#### **B. FIELDS SEARCHED**

Minimum documentation searched (classification system followed by classification symbols)

#### IPC6: C12N, C11B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

## SE,DK,FI,NO classes as above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

## WPI, CA, MEDLINE, BIOSIS, DBA, FSTA, EMBL/PIR/SWISSPROT/GENESEQ

## C. DOCUMENTS CONSIDERED TO BE RELEVANT Category\* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. EP 0513709 A2 (RÖHM GMBH), 19 November 1992 23-25,27 (19.11.92), the claims A 1-22,26 EP 0622446 A2 (SHOWA SANGYO CO., LTD.), A 1-27 2 November 1994 (02.11.94) Biosci. Biotech. Biochem., Volume 60, No 7, 1996, Hideki Oishi et al, "Purification and Some Properties of Phospholipase B from A 1-22 Schizosaccharomyces pombe" page 1087 - page 1092

	Further documents are listed in the continuation of Box	x C.	X See patent family annex.
* 'A' 'E' 'L' 'O' 'P'	Special categories of cited documents:  document defining the general state of the art which is not considered to be of particular relevance ertier document but published on or after the international filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than	"T" "X" "Y"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention  document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone  document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
<u> </u>	the priority date claimed	<b>"</b> &"	document member of the same patent family
Date	e of the actual completion of the international search	Date	of mailing of the international search report
18	February 1998		2 4 -02- 1998
Nan	ne and mailing address of the ISA/	Autho	rized officer
Swe	edish Patent Office		
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